

Document made available under the Patent Cooperation Treaty (PCT)

International application number: PCT/GB05/050016

International filing date: 15 February 2005 (15.02.2005)

Document type: Certified copy of priority document

Document details: Country/Office: GB
Number: 0404420.2
Filing date: 27 February 2004 (27.02.2004)

Date of receipt at the International Bureau: 07 April 2005 (07.04.2005)

Remark: Priority document submitted or transmitted to the International Bureau in compliance with Rule 17.1(a) or (b)



World Intellectual Property Organization (WIPO) - Geneva, Switzerland
Organisation Mondiale de la Propriété Intellectuelle (OMPI) - Genève, Suisse



PCT/GB2005/050016.



INVESTOR IN PEOPLE

The Patent Office
Concept House
Cardiff Road
Newport
South Wales
NP10 8QQ

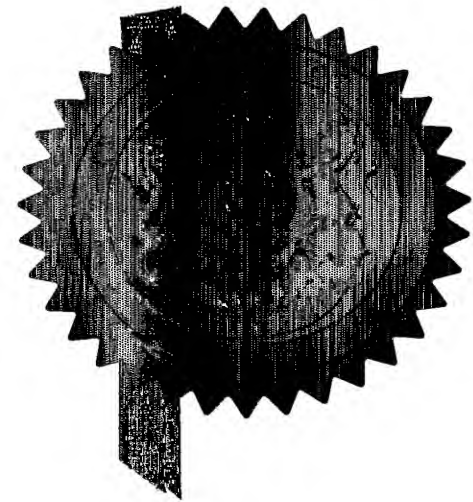
I, the undersigned, being an officer duly authorised in accordance with Section 74(1) and (4) of the Deregulation & Contracting Out Act 1994, to sign and issue certificates on behalf of the Comptroller-General, hereby certify that annexed hereto is a true copy of the documents as originally filed in connection with the patent application identified therein.

I also certify that the application is now proceeding in the name as identified herein.

In accordance with the Patents (Companies Re-registration) Rules 1982, if a company named in this certificate and any accompanying documents has re-registered under the Companies Act 1980 with the same name as that with which it was registered immediately before re-registration save for the substitution as, or inclusion as, the last part of the name of the words "public limited company" or their equivalents in Welsh, references to the name of the company in this certificate and any accompanying documents shall be treated as references to the name with which it is so re-registered.

In accordance with the rules, the words "public limited company" may be replaced by p.l.c., plc, P.L.C. or PLC.

Re-registration under the Companies Act does not constitute a new legal entity but merely subjects the company to certain additional company law rules.



Signed

Andrew Gersey

Dated

9 March 2005





INVESTOR IN PEOPLE

GB 0404420.2

By virtue of a direction given under Section 30 of the Patents Act 1977, the application is proceeding in the name of:

NICHE GENERICS LIMITED,
1 The Cam Centre,
Wilbury Way,
HITCHIN,
Hertfordshire,
SG4 0TW,
United Kingdom

Incorporated in the United Kingdom,

[ADP No. 08774671001]



Request for grant of a patent

27 FEB 2004



1/77

The Patent Office

Cardiff Road
Newport
Gwent NP9 1RH

1. Your reference	ASE/45032GB1		
2. Patent application number	0404420.2		
3. Full name, address and post code of the or each applicant	(1) Niche Generics Limited 1 The Cam Centre Wilbury Way Hitchin Hertfordshire SG4 0DW United Kingdom 08774671001 (2) Rubicon Research Private Limited 221, Annexe Building Goregaon - Mulund Link Road Bhandup (West) Mumbai - 400 078 India 08818759001		
Patents ADP number			
If the applicant is a corporate body, give the country/state of its incorporation	(1) United Kingdom (2) India		
4. Title of the invention	Pharmaceutical Composition		
5. Name of your agent	VENNER, SHIPLEY & CO		
"Address for service" in the United Kingdom to which all correspondence should be sent	20 LITTLE BRITAIN LONDON EC1A 7DH		
Patents ADP	1669004 ✓		
6. If you are declaring priority from one or more earlier patent applications, give the country and the date of filing of the or each of these earlier applications and the or each application number	Country	Priority application number	Date of filing
7. If this application is divided or otherwise derived from an earlier UK application, give the number and filing date of the earlier application	Number of earlier application		Date of Filing

Patents Form 1/77

8. Is a statement of inventorship and of right to grant of a patent required in support of this request? (Answer 'YES' if:
a) any applicant in 3. above is not an inventor, or
b) there is an inventor who is not named as an applicant, or
c) any named applicant is a corporate body)

YES

9. Enter the number of sheets for any of the following items you are filing with this form.
Do not count copies of the same document

Continuation sheets of this form

Description	25	8
Claim(s)	7	
Abstract	1	✓
Drawing(s)	2	

10. If you are also filing any of the following state how many against each item.

Priority documents

Translations of priority documents

Statement of inventorship and right to grant of a patent (*Patents Form 7/77*)

Request for preliminary examination and search (*Patents Form 9/77*)

ONE

Request for substantive examination (*Patents Form 10/77*)

Any other documents

11.

I/We request the grant of a patent on the basis of this application.

Signature

Date

27 February 2004

12. Name and daytime telephone number of person to contact in the United Kingdom

Almut S. Elend
020 7600 4212

Pharmaceutical Composition

Technical field

5 The present invention relates to a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof. In particular, the invention relates to a pharmaceutical composition, which comprises an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, and a C₁₆-C₂₈ glyceride. ACE inhibitors useful in the present invention are susceptible to
10 heat and/or mechanical stress-induced degradation. Preferred ACE inhibitors are ramipril, trandolapril, quinapril and pharmaceutically acceptable salts and derivatives thereof. The composition of the present invention may be for use as a medicament for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension,
15 cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

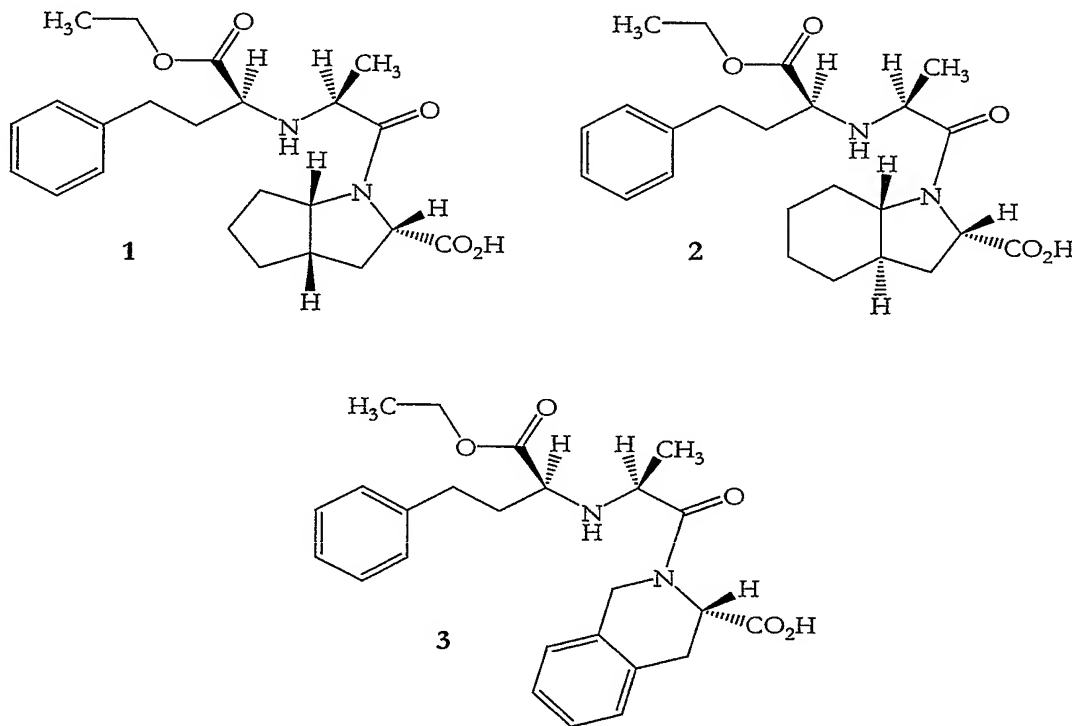
The present invention further relates to a method of preparing the pharmaceutical composition of the present invention. The present invention also relates to a method of providing a stable pharmaceutical composition comprising an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, by incorporating a C₁₆-C₂₈ glyceride into the composition. The present invention further relates to a use of a C₁₆-C₂₈ glyceride to provide a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof.
25

Background art

ACE inhibitors, i.e. inhibitors of angiotensin converting enzymes, are drugs useful in the treatment of cardiovascular disorders, in particular hypertension and coronary
30 heart disease. It has been widely observed that ACE inhibitors are susceptible to degradation between the time of manufacture and the time of desired usage, in particular due to cyclization, hydrolysis and oxidation. Typical degradation products

are hydrolytic degradation products formed by hydrolysis of the ACE inhibitor and diketopiperazine degradation products formed by cyclization of the ACE inhibitor.

Ramipril, also called $(2S,3aS,6aS)$ - $1-[(2S)-[(1S)-(ethoxycarbonyl)-3-$ phenylpropyl]amino]-1-oxopropyl]-octahydrocyclopenta[b]pyrrole-2-carboxylic acid, is an ACE inhibitor of formula 1. Trandolapril, also called $(2S,3aR,7aS)$ - $1-[(2S)-[(1S)-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-octahydro-1H-indole-2-carboxylic acid, is an ACE inhibitor of formula 2. Quinapril, also called $(3S)-2-[(2S)-[(1S)-(ethoxycarbonyl)-3-phenylpropyl]amino]-1-oxopropyl]-1,2,3,4-tetrahydroisoquinoline-3-carboxylic acid, is an ACE inhibitor of formula 3.$$



ACE inhibitors such as ramipril, trandolapril or quinapril, are used in the treatment or prevention of cardiovascular diseases, coronary heart diseases, peripheral vascular diseases, arrhythmias, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

Currently commercially available formulations of ramipril contain as inactive ingredients one or more of the following excipients: hydroxypropylcellulose, hydroxypropylmethylcellulose, microcrystalline cellulose, pregelatinized starch, maize starch, sodium stearyl fumarate, gelatin, anhydrous lactose, polyethylene glycol, polyoxyl hydrogenated castor oil, propyl gallate, sodium aluminium silicate, paraffin, and/or colouring agents (such as black, red and/or yellow ferric oxide E172, titanium dioxide E171, and/or indigo carmine E132).

Currently commercially available formulations of trandolapril contain as inactive ingredients one or more of the following excipients: corn starch, lactose, povidone, and/or sodium stearyl fumarate.

Currently commercially available formulations of quinapril contain as inactive ingredients one or more of the following excipients: magnesium carbonate, lactose, hydrous lactose, gelatin, povidone, crospovidone, magnesium stearate, candelilla wax, hydroxypropylmethylcellulose, hydroxypropylcellulose, polyethylene glycol, maize starch, talc, and/or colouring agents (such as red and/or yellow ferric oxide E172, titanium dioxide E171, and/or indigotine E132).

Many ACE inhibitors, including ramipril, trandolapril and quinapril, have an ester (CO-O) and/or an amide (CO-N) bond. Such bonds are susceptible to hydrolysis leading to the formation of hydrolytic degradation products. Moreover, due to their molecular structure many ACE inhibitors, including ramipril, trandolapril and quinapril, are susceptible to cyclization to form diketopiperazine degradation products. Some known degradation products of ramipril are shown in Figure 1, including hydrolytic degradation products E and F, and diketopiperazine degradation products D, K and L.

The degradation of ACE inhibitors has been found to occur both in solid and in liquid states. As the degradation of an ACE inhibitor in a pharmaceutical composition increases, the concentration of available, functional ACE inhibitor decreases. Thus the shelf-life of pharmaceutical compositions comprising the ACE

inhibitor is limited due to this degradation. Accordingly, degradation should be avoided.

5. Various ways to minimize the degradation of ACE inhibitors in pharmaceutical compositions have been advocated. For example, it has been suggested that alkali or alkaline-earth metal salts can stabilise ACE inhibitors and their salts and derivatives in pharmaceutical compositions.

10 WO 01/15724 and US-6,555,551 disclose a method of stabilising pharmaceutical compositions comprising ACE inhibitors such as ramipril hydrochloride or quinapril hydrochloride. The method comprises the step of mixing an alcoholic dispersion of an ACE inhibitor with an aqueous solution or dispersion of a metal compound; the resulting mixture may be dried. Suitable metal compounds are alkali or alkaline-earth metal salts.

15 EP-0,280,999 and US-4,743,450 teach that the cyclization, hydrolysis and discolouration of pharmaceutical compositions, comprising quinapril, enalapril, indolapril or structurally-related ACE inhibitors, are minimized by formulating the compositions with a metal-containing alkaline stabilizer. The metal-containing 20 alkaline stabilizer is preferably an inorganic salt of an alkali or alkaline-earth metal, such as magnesium, calcium or sodium borate, silicate or carbonate.

25 WO 03/059388 discloses that the cyclization, hydrolysis and discolouration of pharmaceutical compositions, comprising ramipril, quinapril, trandolapril or structurally-related ACE inhibitors, are minimized by formulating the compositions with a basic compound and a filler. The basic compound is preferably an alkali or alkaline-earth metal carbonate, such as magnesium carbonate, sodium carbonate or sodium hydrogen carbonate. The filler is preferably an insoluble alkaline-earth metal hydrogen phosphate, such as calcium hydrogen phosphate.

30 WO 02/11709 discloses stable pharmaceutical compositions comprising ramipril and an effervescent system. The effervescent system comprises an alkali or alkaline-earth metal carbonate or bicarbonate, such as sodium, calcium or magnesium

carbonate or bicarbonate, and at least one acid, such as citric acid, monosodium citrate, ascorbic acid, gluconic acid, lactic acid, malic acid or tartaric acid. The ratio of acid to (bi)carbonate is said to be between 0.6 and 1.3, and the ratio of ramipril to effervescent system is said to be between 0.004 and 0.013, for the pharmaceutical compositions to be stable.

WO 99/62560 and US-6,417,196 disclose pharmaceutical compositions, comprising quinapril, enalapril, indolapril or structurally-related ACE inhibitors, which are stabilised by the presence of magnesium oxide, preferably in combination with a hydrolysis-minimizing agent. The presence of magnesium oxide is also said to lend itself to favourable processing conditions during the manufacture of the ACE inhibitor-containing compositions, especially processing by wet granulation.

It has also been suggested that certain acids can be used to stabilise ACE inhibitors in pharmaceutical compositions. EP-0,468,929, US-6,300,361 and US-6,300,362 disclose the use of hydrochloric acid donors as stabilizers in pharmaceutical compositions comprising ACE inhibitors such as quinapril, enalapril, spirapril, spiraprilate, ramipril, perindopril, indolapril, lisinopril, alacepril, trandolapril, benazepril, libenzapril, delapril or cilazapril. Suitable hydrochloric acid donors are amino acid hydrochlorides, such as glycine, glutamic acid, betaine, alanine, valine, lysine, arginine or aspartic acid hydrochloride, and Lewis acid chlorides, such as ferric, zinc or aluminium chloride.

Furthermore, it has been suggested that certain compounds such as lactose monohydrate can be used to stabilise ACE inhibitors such as ramipril in pharmaceutical compositions. WO 03/028707 discloses pharmaceutical compositions comprising ramipril and lactose monohydrate as diluent. The lactose monohydrate was found to stabilise the ramipril in the compositions. The compositions may further optionally comprise a lubricant, such as magnesium, zinc or calcium stearate.

Moreover, the use of protective coatings has been advocated to stabilise ACE inhibitors in pharmaceutical compositions. EP-0,317,878, US-5,151,433 and US-

5,442,008 disclose pharmaceutical compositions comprising ACE inhibitors such as ramipril, enalapril, perindopril, indolapril, lisinopril, quinapril, alacepril or trandolapril, in which the ACE inhibitors are stabilised by a polymeric protective coating and/or by a buffer which maintains the pH of the compositions between 5.5
5 and 8.0.

WO 95/34283, EP-0,624,364 and US-5,527,540 disclose pharmaceutical compositions comprising an alkali-sensitive active substance, such as captopril, ramipril, perindopril erbumine or enalapril, and an effervescent system, such as a
10 carbonate component. To stabilise the active substance, it is embedded in at least one of the following compounds: an edible organic acid, a higher alcohol, a hydrocolloid, a long-chain polyvinylpyrrolidone, and is preferably coated with at least one of said compounds. The carbonate component is also preferably embedded in at least one edible organic acid and coated by the same or another
15 acid.

Furthermore, in WO 03/059330 it has been suggested that mechanical stress-induced degradation of ACE inhibitors such as ramipril, spirapril, lisinopril, enalapril, quinapril, benazepril or structurally-related ACE inhibitors, can be
20 avoided by coating a core of diluents and other formulating agents with a layer of the ACE inhibitor. The core is compressed prior to coating with the ACE inhibitor, thereby avoiding the need to compress the ACE inhibitor and thus avoiding mechanical stress-induced degradation.

25 It has still further been suggested to stabilise ACE inhibitors by derivatisation. For example, WO 02/03970 discloses a transdermal therapeutic system comprising an adhesive matrix. The matrix comprises a derivative of an ACE inhibitor such as ramipril or trandolapril, which has been stabilised by derivatisation into a salt or diester.

30

Despite these efforts to stabilise ACE inhibitors, there remains a long-standing need for stable pharmaceutical compositions comprising an ACE inhibitor or a

pharmaceutically acceptable salt or derivative thereof, and methods of preparing the same.

Surprisingly, it has now been found that the presence of a C₁₆-C₂₈ glyceride reduces
5 or slows the degradation of ACE inhibitors such as ramipril, trandolapril, quinapril, or their salts or derivatives in pharmaceutical compositions. Astonishingly, until now C₁₆-C₂₈ glycerides such as glycerol dibehenate, a common pharmaceutical excipient, have not been used in pharmaceutical compositions comprising ramipril, trandolapril, quinapril, or their salts or derivatives either in the published prior art
10 or in commercially available compositions.

Summary of the invention

For the purposes of the present invention, a "C₁₆-C₂₈ glyceride" is a mono-, di- or
15 tri-glyceride comprising one, two or three C₁₆-C₂₈ acyl moieties respectively. Preferably each C₁₆-C₂₈ acyl moiety is independently of the formula -CO-R, wherein R is a saturated or unsaturated hydrocarbon, which contains from 16 to 28 carbon atoms, and which is straight-chained or branched. Preferably R is a saturated hydrocarbon. Preferably R is a straight-chained hydrocarbon. The acyl moieties
20 may be derived from naturally occurring or synthetic fatty acids. The terms "C₁₈-C₂₆ glyceride", "C₂₀-C₂₄ glyceride" and "C₂₂ glyceride" are defined accordingly. Glycerol dibehenate comprises mainly C₂₂ diglyceride comprising two C₂₂ acyl moieties of the formula -CO-(CH₂)₂₀-CH₃.

25 A pharmaceutical composition comprising an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, is considered to be "stable", if the ACE inhibitor, or its salt or derivative, in the pharmaceutical composition degrades less or more slowly than it does in known pharmaceutical compositions. The term "unstable" is defined accordingly.

30 An excipient is considered to be "compatible" with an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, if it does not promote the degradation of the ACE inhibitor, or its salt or derivative, i.e. if the ACE inhibitor,

or its salt or derivative, does not degrades more or faster in the presence of the excipient compared to the degradation of the ACE inhibitor, or its salt or derivative, on its own. The terms "compatibility", "incompatible" and "incompatibility" are defined accordingly.

5 An ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, is considered to be "susceptible to heat and/or mechanical stress-induced degradation", if it degrades more or faster when it is subjected to heat and/or mechanical stress such as, for example, due to pressure and heat exerted during
10 compression of a powder blend into tablets, than it does when it is not subjected to heat and/or mechanical stress.

A drug, such as an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof, and an excipient, such as glycerol dibehenate, are considered to form a
15 "mixture", if the drug and the excipient are blended together. Thus, if a first excipient is solely used to coat a drug or a drug/second excipient blend, then the first excipient is not considered to form a mixture with the drug or the drug/second excipient blend. However, if an excipient is blended together with a drug and is also used to coat the drug/excipient blend, then the excipient is considered to form
20 a mixture with the drug. A mixture or blend of a drug and an excipient is considered to form an "intimate mixture or blend", if the mixture or blend is substantially uniform.

25 A first embodiment of the present invention provides a pharmaceutical composition comprising an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, and a C₁₆-C₂₈ glyceride. Preferably the pharmaceutical composition comprises 5-30% by weight C₁₆-C₂₈ glyceride, more preferably 5-20% by weight, even more preferably 10-15% by weight of the total composition.

30 Preferably the glyceride comprises one, two or three C₁₆-C₂₈ acyl moieties, wherein each C₁₆-C₂₈ acyl moiety is independently of the formula -CO-R, wherein R is a saturated or unsaturated hydrocarbon, which contains from 16 to 28 carbon atoms, and which is straight-chained or branched. Preferably R is a saturated hydrocarbon

and/or R is a straight-chained hydrocarbon. Preferably the glyceride is a C₁₈-C₂₆ glyceride, more preferably a C₂₀-C₂₄ glyceride, even more preferably a C₂₂ glyceride. Preferably the glyceride comprises at least 50% diglyceride, more preferably at least 60% diglyceride, even more preferably at least 70% diglyceride. In the most preferred embodiment of the present invention, the glyceride is glycerol dibehenate.

Preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is susceptible to heat and/or mechanical stress-induced degradation. More preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril, trandolapril, quinapril, or a pharmaceutically acceptable salt or derivative thereof. In the most preferred embodiment of the present invention, the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril or a pharmaceutically acceptable salt or derivative thereof.

Preferably the pharmaceutical composition comprises one or more further excipients, which are compatible with the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.

The one or more further excipients may be selected from carbonates (such as calcium carbonate, sodium carbonate or magnesium carbonate), phosphates (such as anhydrous dibasic calcium phosphate, dibasic calcium phosphate dihydrate, tribasic calcium phosphate or sodium phosphate), sulfates (such as calcium sulfate), silicates (such as kaolin, talc, sodium aluminium silicate, magnesium aluminium silicate, magnesium silicate or magnesium trisilicate), carbohydrates (such as dextrose, dextrin, maltodextrin, dextrose, polydextrose, fructose, sucrose, sugar spheres, compressible sugar, confectioner's sugar, maltose, mannitol, lactose, anhydrous lactose, hydrous lactose, lactitol, maltitol, sorbitol, sodium alginate, alginic acid or liquid glucose), starches (such as starch, pregelatinized starch, maize starch, corn starch or sodium starch glycolate), celluloses (such as carboxymethylcellulose calcium, carboxymethylcellulose sodium, cross-linked carboxymethylcellulose sodium, microcrystalline cellulose, silicified microcrystalline cellulose, powdered cellulose, cellulose acetate, cellulose acetate phthalate, methylcellulose, ethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, low-substituted

hydroxypropylcellulose or hydroxypropylmethylcellulose), polyvinylpyrrolidones (such as povidone or crospovidone), fatty acids or fatty acid derivatives (such as hydrogenated vegetable oil, hydrogenated castor oil, polyoxyl hydrogenated castor oil, mineral oil, light mineral oil, cottonseed oil, a medium-chain triglyceride,
5 glyceryl palmitostearate, calcium stearate, stearic acid, glyceryl monostearate, magnesium stearate, polyoxyethylene stearate, zinc stearate, sodium stearyl fumarate, candelilla wax or glycerol dibehenate), gums (such as tragacanth gum, guar gum or acacia), colouring agents (such as black, red or yellow ferric oxide, titanium dioxide or indigotine), magnesium oxide, sodium chloride, polymethacrylate, propyl
10 gallate, colloidal silicon dioxide, polacrilin potassium, sodium lauryl sulfate, a poloxamer, polyethylene glycol, sodium benzoate, a carbomer, ceratonia, gelatin, paraffin, polyethylene oxide, zein, or a mixture thereof.

Preferably, the one or more further excipients are selected from carbonates (preferably magnesium carbonate), phosphates (preferably anhydrous dibasic calcium phosphate, dibasic calcium phosphate dihydrate or tribasic calcium phosphate), silicates (preferably kaolin, talc, sodium aluminium silicate, magnesium aluminium silicate, magnesium silicate or magnesium trisilicate), carbohydrates (preferably dextrates, maltodextrin, dextrose, polydextrose, fructose, sucrose, sugar
15 spheres, compressible sugar, confectioner's sugar, maltose, mannitol, lactose, anhydrous lactose, hydrous lactose, lactitol, maltitol, sorbitol or sodium alginate), starches (preferably starch, pregelatinized starch, maize starch, corn starch or sodium starch glycolate), celluloses (preferably carboxymethylcellulose calcium, carboxymethylcellulose sodium, cross-linked carboxymethylcellulose sodium,
20 microcrystalline cellulose, powdered cellulose, cellulose acetate, cellulose acetate phthalate, methylcellulose, ethylcellulose, hydroxyethylcellulose, hydroxypropylcellulose, low-substituted hydroxypropylcellulose or hydroxypropylmethylcellulose), polyvinylpyrrolidones (preferably povidone or crospovidone), fatty acids or fatty acid derivatives (preferably hydrogenated vegetable oil, hydrogenated castor oil, polyoxyl hydrogenated castor oil, glyceryl palmitostearate, calcium stearate, stearic acid, glyceryl monostearate, magnesium
25 stearate, zinc stearate, sodium stearyl fumarate, candelilla wax or glycerol dibehenate), gums (preferably guar gum), colouring agents (preferably black, red or

yellow ferric oxide, titanium dioxide or indigotine), sodium chloride, polymethacrylate, propyl gallate, colloidal silicon dioxide, sodium lauryl sulfate, a poloxamer, polyethylene glycol, sodium benzoate, a carbomer, ceratonia, gelatin, paraffin, polyethylene oxide, zein, or a mixture thereof.

5

More preferably, the one or more further excipients are selected from hydroxypropylmethylcellulose, pregelatinised starch, microcrystalline cellulose, lactose, sodium starch glycolate, sodium stearyl fumarate, red ferric oxide and yellow ferric oxide.

10

Preferably the pharmaceutical composition comprises:

1-8% by weight ACE inhibitor, preferably 2-6% by weight;

5-20% by weight C₁₆-C₂₈ glyceride, preferably 10-15% by weight;

60-80% by weight lactose anhydrous, preferably 65-75% by weight;

15

5-20% by weight sodium starch glycolate, preferably 10-15% by weight;

0.5-4 by weight sodium stearyl fumarate, preferably 0.5-2% by weight;

0-0.4% by weight yellow ferric oxide; and

0-0.1% by weight red ferric oxide.

20

Preferably the pharmaceutical composition of the present invention is stable.

Preferably the pharmaceutical composition of the present invention is suitable for direct compression into tablets.

25

Preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and the C₁₆-C₂₈ glyceride form a mixture, preferably an intimate mixture, in the pharmaceutical composition of the present invention. If one or more further excipients are present in the composition, preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, the C₁₆-C₂₈ glyceride and one or more of the one or more further excipients form a mixture, preferably an intimate mixture, in the pharmaceutical composition. Preferably the mixture or the intimate mixture is suitable for direct compression into tablets.

30

5 Optionally the pharmaceutical composition of the present invention comprises granules or particles comprising the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, wherein the granules or particles comprise a coating comprising the C₁₆-C₂₈ glyceride. The granules or particles may optionally further comprise one or more excipients...

Preferably the composition is a solid composition, more preferably it is a non-effervescent composition.

10 Optionally the pharmaceutical composition of the present invention may further comprise a β-blocker, a diuretic, a calcium-channel blocker, a vasodilator anti-hypertensive drug, or an angiotensin II receptor antagonist.

15 Typically, the pharmaceutical composition of the present invention is suitable for oral, parental, transdermal, airway, rectal, vaginal or topical administration. Preferably the composition is suitable for oral administration.

20 A composition suitable for oral administration may be in unit dosage form comprising 1-20mg, preferably 1-10mg, of the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof. A composition suitable for oral administration is typically provided in the form of tablets, capsules, caplets, troches, lozenges, dragées, powder, granules or particles. Optionally the tablets, capsules, caplets, troches, lozenges, dragées, powder, granules or particles not only contain the C₁₆-C₂₈ glyceride, but also comprise a coating comprising the C₁₆-C₂₈ glyceride. Preferably the composition is provided in the form of tablets. Preferably the tablets have a disintegration time of not more than 10 minutes, more preferably of not more than 5 minutes, in water at 36-38°C. Preferably the tablets have a shelf-life of at least 18 months, preferably of at least 24 months, more preferably of at least 4 or 5 years.

25

30 Preferably the composition is for use as a medicament, typically for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular

disease, a peripheral vascular disease, arrhythmia, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

A further embodiment of the present invention provides a method of treating or
5 preventing a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or angina, comprising administering an effective amount of a pharmaceutical composition of the present invention to a patient in need thereof.

10

A further embodiment of the present invention provides a use of a pharmaceutical composition of the present invention in the manufacture of a medicament for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension, 15 cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

A further embodiment of the present invention provides a method of preparing a pharmaceutical composition of the present invention, comprising the step of blending the ACE inhibitor, or the pharmaceutically acceptable salt or derivative 20 thereof, with the C₁₆-C₂₈ glyceride and optionally one or more further excipients. Preferably the method comprises the steps of blending the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, with the C₁₆-C₂₈ glyceride to form a pre-mix, and then blending the pre-mix with one or more further excipients. Preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative 25 thereof, the C₁₆-C₂₈ glyceride and optionally one or more further excipients are blended to form an intimate mixture. Preferably the method further comprises the step of compressing the blend of the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and the excipient(s) into tablets by direct compression. Optionally the tablets, comprising a C₁₆-C₂₈ glyceride, may also be 30 provided with a coating comprising a C₁₆-C₂₈ glyceride.

Alternatively, a method of preparing a pharmaceutical composition of the present invention may comprise the steps of preparing granules or particles comprising the

ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and optionally one or more excipients, and providing the granules or particles with a coating comprising the C₁₆-C₂₈ glyceride.

5 The composition may be prepared by the methods of the present invention in batches of 5-150kg, preferably in batches of 5-100kg.

A further embodiment of the present invention provides a method of providing a stable pharmaceutical composition comprising an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, the method comprising incorporating a C₁₆-C₂₈ glyceride into the composition. Preferably the method of providing a stable pharmaceutical composition comprises incorporating the C₁₆-C₂₈ glyceride into the composition in a mixture, preferably an intimate mixture, with the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.
10 Preferably the pharmaceutical composition is stabilised to minimize the degradation of the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof. Preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is susceptible to heat and/or mechanical stress-induced degradation. More preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril, trandolapril, quinapril, or a pharmaceutically acceptable salt or derivative thereof. In the most preferred embodiment of the present invention, the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril or a pharmaceutically acceptable salt or derivative thereof. Preferably the pharmaceutical composition comprises 5-30% by weight C₁₆-C₂₈ glyceride, more
15 preferably 5-20% by weight, even more preferably 10-15% by weight of the total composition. Preferably the glyceride is a C₁₈-C₂₆ glyceride, more preferably a C₂₀-C₂₄ glyceride, even more preferably a C₂₂ glyceride. Preferably the glyceride comprises at least 50% diglyceride, more preferably at least 60% diglyceride, even more preferably at least 70% diglyceride. In the most preferred embodiment of the
20 present invention, the glyceride is glycerol dibehenate.
25
30

The present invention further provides a use of a C₁₆-C₂₈ glyceride to provide a stable pharmaceutical composition comprising an ACE inhibitor or a

pharmaceutically acceptable salt or derivative thereof. Preferably the pharmaceutical composition is stabilised to minimize the degradation of the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof. Preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is susceptible to heat and/or mechanical stress-induced degradation. More preferably the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril, trandolapril, quinapril, or a pharmaceutically acceptable salt or derivative thereof. In the most preferred embodiment of the present invention, the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril or a pharmaceutically acceptable salt or derivative thereof. Preferably the pharmaceutical composition comprises 5-30% by weight C₁₆-C₂₈ glyceride, more preferably 5-20% by weight, even more preferably 10-15% by weight of the total composition. Preferably the glyceride is a C₁₈-C₂₆ glyceride, more preferably a C₂₀-C₂₄ glyceride, even more preferably a C₂₂ glyceride. Preferably the glyceride comprises at least 50% diglyceride, more preferably at least 60% diglyceride, even more preferably at least 70% diglyceride. In the most preferred embodiment of the present invention, the glyceride is glycerol dibehenate.

Brief description of the drawings

The present invention will now be described by way of example with reference to the accompanying drawings in which:

Figure 1 schematically depicts some degradation pathways of ramipril, in particular to degradation products D, E, F, K and L. A list of degradation products and impurities A to N of ramipril is provided in *European Pharmacopoeia*, 2002, 4th edition.

Figure 2 is a graph showing the increase in total impurities (%) in tablets of formulations 1-4 and 19 when stored at 40°C and 75% relative humidity.

Detailed description of the invention

It has now surprisingly been found that the presence of a C₁₆-C₂₈ glyceride, such as glycerol dibehenate, reduces or slows the degradation of certain ACE inhibitors in pharmaceutical compositions.

5 Glycerol dibehenate, also called glyceryl dibehenate, glyceryl behenate and 2,3-dihydroxypropyl docosanoate, is sold under the trade name Compritol®. PharmEuropa (section 9, 2001) describes glyceryl dibehenate as a mixture of diacylglycerols, mainly dibehenoyleglycerol, together with variable quantities of mono- and triacylglycerols. The US Pharmacopeia 24 / National Formulary 19
10 describes glyceryl behenate as a mixture of glycerides of fatty acids, mainly behenic acid and specifies that the content of 1-monoglycerides should be between 12.0-18.0%.

15 Glycerol dibehenate is used in cosmetics, foods and oral pharmaceutical formulations and is generally regarded as a relatively non-irritant and non-toxic material. It is GRAS listed and included in the FDA's Inactive Ingredients Guide.

20 In pharmaceutical formulations, glycerol dibehenate is mainly used as a tablet or capsule lubricant, tablet binder or coating agent. It has also been investigated for use in the preparation of sustained release tablets.

Without wishing to be bound by theory, it is believed that C₁₆-C₂₈ glycerides, such as glycerol dibehenate, may reduce or slow the degradation of certain ACE inhibitors and their salts and derivatives in pharmaceutical compositions as follows.

25 Certain ACE inhibitors and their salts and derivatives are unstable and susceptible to degradation, especially in the presence of heat and mechanical stress such as, for example, due to pressure and heat exerted during compression of a powder blend into tablets. C₁₆-C₂₈ glycerides, such as glycerol dibehenate, are plastically deformable compounds. Therefore, when pharmaceutical compositions comprising such an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, and a C₁₆-C₂₈ glyceride, are compressed into tablets, the C₁₆-C₂₈ glyceride reduces the compression heat and mechanical stress due to its plastic deformability. In

other words, the C₁₆-C₂₈ glyceride acts as a cushioning agent to protect the unstable ACE inhibitor, or its salt or derivative, from heat and mechanical stress.

Compatibility studies

5

Example 1

Commercially available formulations of ramipril are currently sold by Aventis, Hoechst and Astra under trade names such as Tritace®, Acovil®, Delix® or Ramace®.

10 These commercially available formulations contain ramipril as active ingredient as well as hydroxypropylmethylcellulose, pregelatinised starch, microcrystalline cellulose, sodium stearyl fumarate, yellow ferric oxide and red ferric oxide as inactive ingredients.

15 Ramipril tablets of formulations 1 to 4 were prepared with a composition similar to these commercially available ramipril formulations. Ramipril tablets 1 to 4 all comprise ramipril as well as pregelatinised starch, microcrystalline cellulose, sodium stearyl fumarate and yellow ferric oxide as inactive ingredients, as summarised in Table 1. Ramipril tablets 1 to 4 were prepared by mixing ramipril and the excipients 20 intimately and then compressing the drug/excipient blend into tablets.

Ingredients (mg/tablet)	Tablet 1	Tablet 2	Tablet 3	Tablet 4
Ramipril	2.5	5.0	2.5	5.0
Pregelatinised starch	30.9	61.8	45.9	91.8
Microcrystalline cellulose	65.0	130.0	50.0	100.0
Sodium stearyl fumarate	1.0	2.0	1.0	2.0
Yellow ferric oxide	0.6	1.2	0.6	1.2
Total weight	100.0	200.0	100.0	200.0

Table 1

25 The stability of ramipril in the tablets of formulations 1 to 4 stored in PVdC-coated PVC/aluminium blister packs at 25°C and 60% relative humidity, at 30°C and 60% relative humidity, and at 40°C and 75% relative humidity was studied following the

procedures described in the ICH Guidelines (International Conference on Harmonisation of Technical Standards Guidelines).

The results of the stability studies of ramipril tablets 1 to 4 are presented in Table 2.

- 5 As can bee seen, although immediately after compression of the tablets 1 to 4 there is no significant increase in total impurities, after 6 weeks storage at 40°C and 75% relative humidity, the total impurities in all of tablets 1 to 4 have increased to more than 2.5%. This level of degradation is high and stabilisation of the drug is desirable.

Tablet	Storage conditions	Time	Total Known Impurities (%)	Total Unknown Impurities (%)	Total Impurities (%)
Tablet 1	Initial	Initial	0.64	0.00	0.64
	25°C/60%RH	After 2 weeks	0.84	0.00	0.84
	40°C/75%RH		2.21	0.00	2.21
	25°C/60%RH	After 4 weeks	1.03	0.00	1.03
	40°C/75%RH		3.00	0.07	3.07
	25°C/60%RH	After 6 weeks	1.36	0.00	1.36
	30°C/60%RH		1.67	0.00	1.67
	40°C/75%RH		3.36	0.18	3.54
Tablet 2	Initial	Initial	0.54	0.00	0.54
	25°C/60%RH	After 2 weeks	1.06	0.00	1.06
	40°C/75%RH		2.04	0.00	2.04
	25°C/60%RH	After 4 weeks	1.05	0.06	1.11
	40°C/75%RH		3.43	0.06	3.49
	25°C/60%RH	After 6 weeks	1.53	0.00	1.53
	30°C/60%RH		1.78	0.00	1.78
	40°C/75%RH		3.27	0.25	3.52
Tablet 3	Initial	Initial	0.48	0.00	0.48
	25°C/60%RH	After 2 weeks	0.48	0.00	0.48
	40°C/75%RH		0.96	0.00	0.96
	25°C/60%RH	After 4 weeks	0.85	0.00	0.85
	30°C/60%RH		1.09	0.00	1.09
	40°C/75%RH		2.30	0.05	2.35
	25°C/60%RH	After 6 weeks	0.85	0.00	0.85
	30°C/60%RH		1.08	0.00	1.08
Tablet 4	Initial	Initial	0.56	0.00	0.56
	25°C/60%RH	After 2 weeks	0.98	0.00	0.98
	40°C/75%RH		1.66	0.00	1.66
	25°C/60%RH	After 4 weeks	0.85	0.00	0.85
	30°C/60%RH		1.07	0.06	1.12
	40°C/75%RH		2.25	0.05	1.12
	25°C/60%RH	After 6 weeks	0.93	0.00	0.93
	30°C/60%RH		1.14	0.05	1.19
	40°C/75%RH		2.67	0.07	2.74

Table 2

Example 2

Ramipril tablets of formulations 5 to 18, comprising ramipril and excipients as set out in Table 3, were prepared and the effect of heat and mechanical stress on drug stability in these tablets was studied in order to identify excipients that have a stabilising effect on ramipril. Unless otherwise indicated in Table 3, ramipril tablets 5 to 18 were prepared by mixing ramipril and the excipients intimately and then compressing the drug/excipient blend into tablets.

The stability of ramipril in tablets 5 to 18 stored in high-density polyethylene containers at 40°C and 75% relative humidity was studied following the procedures described in the ICH Guidelines. The results of the stability studies of ramipril tablets 5 to 18 are presented in Table 4.

Based on the results presented in Table 4, it can be concluded that the addition of pH modulators like sodium bicarbonate, lysine monohydrate, magnesium carbonate etc. can help in controlling levels of impurity D, the major heat degradation product of ramipril. However, all of these pH modulators cause a significant increase in impurities E and F, hydrolytic degradation products of ramipril, which are known to occur in alkaline conditions.

Two formulations, tablets 6 and 14, showed a lesser amount of ramipril degradation compared to all other formulations. In both cases, heat degradation product D was found to be less than 1.5% after 8 weeks storage at 40°C and 75% relative humidity, whereas it was almost 6% for formulations 7 and 16. A disadvantage of formulation 14 is that the amount of hydrolytic degradation products E and F increases to almost 1% after 8 weeks storage at 40°C and 75% relative humidity, due to the alkaline pH of formulation of 14. Formulation 6, on the other hand, does not show any hydrolytic degradation products E or F, and the levels of heat degradation product D are also very low.

Ingredients (mg/tablet)	T 5	T 6	T 7	T 8 ^b	T 9 ^b	T 10 ^b	T 11	T 12 ^c	T 13 ^c	T 14 ^{b,d}	T 15 ^b	T 16 ^e	T 17	T 18
Ramipril	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0	5.0
Anhydrous lactose	179.0	164.0	189.0	164.0	-	-	-	-	-	184.0	184.0	149.0	139.0	-
Sodium starch glycolate	4.0	4.0	4.0	4.0	-	-	-	-	-	4.0	4.0	-	4.0	-
Sodium stearyl fumarate	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0	2.0
Stearic acid	10.0	-	-	-	-	-	-	-	-	-	-	-	-	-
Glycerol dibehenate	-	25.0	-	-	-	-	-	-	-	-	-	-	-	-
Lysine monohydrate	-	-	-	25.0	-	-	-	-	-	-	-	-	-	-
Sodium bicarbonate	-	-	-	-	5.0	50.0	-	-	-	-	-	-	-	-
Tricalcium phosphate	-	-	-	-	184.0	139.0	189.0	-	-	-	-	-	-	-
Crospovidone	-	-	-	-	4.0	4.0	4.0	-	-	-	-	-	-	-
Alcoholic solution of polyvinyl pyrrolidone	-	-	-	-	-	-	-	10.0	-	-	-	-	-	-
Alcoholic solution of Carbopol®	-	-	-	-	-	-	-	-	10.0	-	-	-	-	-
Microcrystalline cellulose	-	-	-	-	-	-	-	115.0	115.0	-	-	44.0	-	123.94
Pregelled starch	-	-	-	-	-	-	-	-	-	68.0	68.0	-	-	68.3
TRIS buffer	-	-	-	-	-	-	-	-	-	5.0	5.0	-	-	-
Magnesium carbonate	-	-	-	-	-	-	-	-	-	-	-	50.0	-	-
Hydroxypropylmethylcellulose	-	-	-	-	-	-	-	-	-	-	-	-	0.76	-
Total weight	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0	200.0

a - Abbreviations used: T = tablet.

b - The pH of formulations 8, 9, 10, 14 and 15 was 9.4, 6.2, 8.3, 8.4 and 8.4 respectively.

c - The drug was granulated with the alcoholic solution, the granules were dried, intimately mixed with the other excipients and compressed into tablets 12 and 13.

d - Formulation 14 was prepared by wet granulation.

e - Low moisture excipients were used to control the moisture content of formulation 16.

Table 3

Tab.	Composition	Initial				After 2 weeks				After 4 weeks				After 6 weeks				After 8 weeks			
		Imp. D	Imp. E+F	Total Imp.	Imp. D	Imp. E+F	Total Imp.	Imp. D	Imp. E+F	Total Imp.	Imp. D	Imp. E+F	Total Imp.	Imp. D	Imp. E+F	Total Imp.	Imp. D	Imp. E+F	Total Imp.		
5	ramipril, stearic acid (1:2), AL, SSG, SSF	0.86	0.03	1.10	0.83	0.03	1.85	36.23	0.14	38.38	-	-	-	-	-	-	-	-	-		
6	ramipril, glycerol dibehenate (1:5), AL, SSG, SSF	0.12	0.02	0.25	0.01	0.46	0.39	0.02	0.55	0.70	0.02	0.98	1.25	0.00	1.76	-	-	-	-		
7	ramipril, AL, SSG, SSF (NC)	0.19	0.02	0.35	1.77	0.09	2.09	3.72	0.15	4.15	4.60	0.19	5.35	5.98	0.22	6.82	-	-	-		
8	ramipril, lysine monohydrate (1:5), AL, SSG, SSF	0.08	0.10	0.31	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-		
9	ramipril, sodium bicarbonate (1:1), TP, CP, SSF	0.09	0.28	0.61	0.18	4.13	4.54	-	-	-	-	-	-	-	-	-	-	-	-		
10	ramipril, sodium bicarbonate (1:10), TP, CP, SSF	0.07	0.33	0.64	0.11	2.17	2.43	-	-	-	-	-	-	-	-	-	-	-	-		
11	ramipril, TP, CP, SSF (NC)	0.12	0.24	0.64	0.20	4.51	5.05	-	-	-	-	-	-	-	-	-	-	-	-		
12	ramipril, PVP (1:2), MCC, PS, SSF	0.62	0.05	0.79	15.36	0.53	16.38	-	-	-	-	-	-	-	-	-	-	-	-		
13	ramipril, Carbopol® (1:2), MCC, PS, SSF	0.09	0.05	0.23	0.43	0.60	1.14	0.95	1.69	2.88	1.22	2.39	3.95	-	-	-	-	-	-		
14	ramipril, TRIS buffer (1:1), AL, SSG, SSF (wet granulation)	0.00	0.03	0.08	0.07	0.11	0.25	0.29	0.41	1.00	0.49	0.69	1.53	0.85	0.98	2.47	-	-	-		
15	ramipril, TRIS buffer (1:1), AL, SSG, SSF (dry mix)	0.09	0.02	0.23	0.11	0.16	0.37	0.65	0.71	1.94	1.11	1.37	3.39	-	-	-	-	-	-		
16	ramipril, AL, MCC, SSF (LM)	0.10	0.02	0.26	1.57	0.08	1.73	3.38	0.15	3.74	4.95	0.21	5.50	5.58	0.24	6.20	-	-	-		
17	ramipril, magnesium carbonate (1:10), AL, SSG, SSF	0.05	0.09	0.27	0.11	0.73	0.96	0.18	2.58	2.99	0.21	3.07	3.59	-	-	-	-	-	-		
18	ramipril, HPMC (1:0.76), MCC, PS, SSF	0.11	0.02	0.30	1.41	0.07	1.59	2.67	0.11	2.97	2.72	0.09	3.21	-	-	-	-	-	-		

Abbreviations used: AL = anhydrous lactose; SSG = sodium starch glycolate; SSF = sodium stearyl fumarate; TP = tricalcium phosphate; CP = crospovidone; PVP = polyvinylpyrrolidone; MCC = microcrystalline cellulose; PS = pregelled starch; HPMC = hydroxypropylmethylcellulose; NC = negative control; LM = low moisture; Imp. = impurity; Tab. = tablet.

Table 4

Indeed formulation 6, comprising glycerol dibehenate, was found to be more stable than any of the other formulations, showing no hydrolytic degradation to impurities E and F, and only minimal heat degradation to impurity D. Since the only difference between formulation 6 and less stable formulation 7 is the presence of 5 glycerol dibehenate in formulation 6, it can be concluded that glycerol dibehenate has a stabilising effect on ramipril.

Example 3

10 To confirm the stabilising effect of glycerol dibehenate, ramipril tablets of formulations 19 to 23, comprising ramipril, glycerol dibehenate and other excipients as summarised in Table 5, were prepared and the effect of heat and mechanical stress on drug stability in these tablets was studied. Ramipril tablets 19 to 23 were prepared by pre-mixing ramipril and glycerol dibehenate intimately, followed by 15 mixing the ramipril/glycerol dibehenate pre-mix with the remaining excipients intimately, and then compressing the drug/excipient blend into tablets.

Ingredients (mg/tablet)	Tablet 19	Tablet 20	Tablet 21	Tablet 22	Tablet 23
Ramipril	5.0	10.0	5.0	2.5	1.25
Compritol 888 ATO®	25.0	25.0	25.0	12.5	6.25
Pharmatose DCL 21®	144.0	139.0	143.8	71.6	36.0
Primojel®	24.0	24.0	24.0	12.0	6.0
PRUV®	2.0	2.0	2.0	1.0	0.5
Red ferric oxide	-	-	0.2	-	-
Yellow ferric oxide	-	-	-	0.4	-
Total weight	200.0	200.0	200.0	100.0	50.0

Table 5

20 Compritol 888 ATO® is glycerol dibehenate; Pharmatose DCL 21® is anhydrous lactose; Primojel® is sodium starch glycolate; PRUV® is sodium stearyl fumarate; and red ferric oxide and yellow ferric oxide are colouring agents.

The stability of ramipril in tablet 19 stored in high-density polyethylene containers at 25°C and 60% relative humidity, at 35°C and 60% relative humidity, and at 40°C and 75% relative humidity was studied following the procedures described in the ICH Guidelines. The results of the stability studies of ramipril tablet 19 are
5 presented in Table 6.

Storage conditions	Time	Total Known Impurities (%)	Total Unknown Impurities (%)	Total Impurities (%)
Initial	Initial	0.30	0.00	0.30
25°C/60%RH	After 2 weeks	0.34	0.00	0.34
40°C/75%RH		0.53	0.00	0.53
25°C/60%RH	After 4 weeks	0.39	0.00	0.39
40°C/75%RH		0.60	0.06	0.66
25°C/60%RH	After 6 weeks	0.39	0.00	0.39
40°C/75%RH		0.88	0.09	0.97
25°C/60%RH	After 8 weeks	0.47	0.00	0.47
30°C/60%RH		0.52	0.00	0.52
40°C/75%RH		1.28	0.16	1.44
25°C/60%RH	After 12 weeks	0.48	0.00	0.48
30°C/60%RH		0.55	0.00	0.55
40°C/75%RH		1.95	0.40	2.35

Table 6

The stability of ramipril in tablets 1-4 and 19 is compared in Figure 2, which shows
10 the increase in total impurities (%) in the tablets when stored at 40°C and 75% relative humidity. As can be seen, ramipril in tablet 19 is much more stable to degradation than ramipril in tablets 1-4, which have a composition similar to currently commercially available ramipril formulations.

15 The stability of ramipril in tablets 20 to 23 stored in high-density polyethylene containers at 40°C and 75% relative humidity was studied following the procedures described in the ICH Guidelines. The percentage increase of heat degradation product D is summarised in Table 7.

Time	Impurity D (%)			
	Tablet 20	Tablet 21	Tablet 22	Tablet 23
Initial	0.24	0.30	0.28	0.30
After 2 weeks	0.42	0.32	0.27	0.40
After 4 weeks	0.51	0.67	0.35	0.59
After 6 weeks	0.72	0.95	0.44	0.86
After 8 weeks	0.87	0.95	0.42	1.03
After 12 weeks	1.15	1.68	0.59	1.60

Table 7

The results presented in Tables 6 and 7 confirm that glycerol dibehenate reduces the degradation of ramipril in pharmaceutical compositions.

5

It will be understood that the present invention has been described above by way of example only. The examples are not intended to limit the scope of the invention. Various modifications and embodiments can be made without departing from the scope and spirit of the invention, which is defined by the following claims only.

10

Claims

1. A pharmaceutical composition comprising an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, and a C₁₆-C₂₈ glyceride.

5

2. A pharmaceutical composition of claim 1, comprising 5-30% by weight C₁₆-C₂₈ glyceride.

10 3. A pharmaceutical composition of claim 1 or 2, comprising one or more further excipients which are compatible with the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.

15 4. A pharmaceutical composition of claim 3, wherein the one or more further excipients are selected from hydroxypropylmethylcellulose, pregelatinised starch, microcrystalline cellulose, lactose, sodium starch glycolate, sodium stearyl fumarate, red ferric oxide and yellow ferric oxide.

5. A pharmaceutical composition of any one of the preceding claims, comprising:

20

2-6% by weight ACE inhibitor,

10-15% by weight C₁₆-C₂₈ glyceride,

65-75% by weight lactose anhydrous,

10-15% by weight sodium starch glycolate,

0.5-2% by weight sodium stearyl fumarate,

25 0-0.4% by weight yellow ferric oxide, and

0-0.1% by weight red ferric oxide.

6. A pharmaceutical composition of any one of the preceding claims, wherein the pharmaceutical composition is stable.

30

7. A pharmaceutical composition of any one of the preceding claims, suitable for direct compression into tablets.

8. A pharmaceutical composition of any one of the preceding claims, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and the C₁₆-C₂₈ glyceride form a mixture.
- 5 9. A pharmaceutical composition of any one of the preceding claims, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, the C₁₆-C₂₈ glyceride and one or more further excipients form a mixture.
- 10 10. A pharmaceutical composition of claim 8 or 9, wherein the mixture is an intimate mixture.
11. A pharmaceutical composition of any one of claims 8 to 10, wherein the mixture is suitable for direct compression into tablets.
- 15 12. A pharmaceutical composition of any one of claims 1 to 7, comprising granules or particles comprising the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, wherein the granules or particles comprise a coating comprising the C₁₆-C₂₈ glyceride.
- 20 13. A pharmaceutical composition of any one of the preceding claims, further comprising a β-blocker, a diuretic, a calcium-channel blocker, a vasodilator anti-hypertensive drug, or an angiotensin II receptor antagonist.
- 25 14. A pharmaceutical composition of any one of the preceding claims, wherein the composition is suitable for oral, parental, transdermal, airway, rectal, vaginal or topical administration.
15. A pharmaceutical composition of any one of the preceding claims, wherein the composition is suitable for oral administration.
- 30 16. A pharmaceutical composition of claim 15, wherein the composition is in unit dosage form comprising 1-20mg of the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.

17. A pharmaceutical composition of claim 15 or 16, wherein the composition is provided in the form of a tablet, capsule, caplet, troche, lozenge, dragée, powder, granules or particles.
- 5
18. A pharmaceutical composition of claim 17, wherein the tablet, capsule, caplet, troche, lozenge, dragée, powder, granules or particles comprise a coating comprising the C₁₆-C₂₈ glyceride.
- 10 19. A pharmaceutical composition of any one of claims 15 to 18, wherein the composition is provided in the form of a tablet.
20. A pharmaceutical composition of claim 19, wherein the tablet has a disintegration time of not more than 10 minutes.
- 15
21. A pharmaceutical composition of claim 19 or 20, wherein the tablet has a shelf-life of at least 5 years.
22. A pharmaceutical composition of any one of the preceding claims, substantially as hereinbefore described with reference to the description.
- 20
23. A pharmaceutical composition of any one of the preceding claims, for use as a medicament.
- 25
24. A pharmaceutical composition of claim 23, for use as a medicament for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.
- 30 25. A method of treating or preventing a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or

angina, comprising administering an effective amount of a pharmaceutical composition of any one of claims 1 to 24 to a patient in need thereof.

26. Use of a pharmaceutical composition of any one of claims 1 to 24 in the

5 manufacture of a medicament for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension, cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

10 27. A method of preparing a pharmaceutical composition of any one of claims 1 to 24, comprising the step of blending the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, with the C₁₆-C₂₈ glyceride.

15 28. A method of claim 27, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and the C₁₆-C₂₈ glyceride are blended to form an intimate mixture.

20 29. A method of claim 27 or 28, comprising the step of blending the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, with the C₁₆-C₂₈ glyceride and one or more further excipients.

25 30. A method of claim 29, comprising the steps of blending the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, with the C₁₆-C₂₈ glyceride to form a pre-mix, and blending the pre-mix with one or more further excipients.

31. A method of claim 29 or 30, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, the C₁₆-C₂₈ glyceride and one or more further excipients are blended to form an intimate mixture.

30

32. A method of any one of claims 27 to 31, further comprising the step of compressing the blend of the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and the excipient(s) into tablets by direct compression.

33. A method of claim 32, further comprising the step of providing the tablets with a coating comprising the C₁₆-C₂₈ glyceride.
- 5 34. A method of preparing a pharmaceutical composition of any one of claims 1 to 24, comprising the steps of preparing granules or particles comprising the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, and optionally one or more excipients, and providing the granules or particles with a coating comprising the C₁₆-C₂₈ glyceride.
- 10 35. A method of any one of claims 27 to 34, wherein the pharmaceutical composition is prepared in batches of 5-150kg.
- 15 36. A method of providing a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof, the method comprising incorporating a C₁₆-C₂₈ glyceride into the composition.
- 20 37. A method of claim 36, comprising incorporating the C₁₆-C₂₈ glyceride into the composition in a mixture with the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.
- 25 38. A method of claim 37, comprising incorporating the C₁₆-C₂₈ glyceride into the composition in an intimate mixture with the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.
39. Use of a C₁₆-C₂₈ glyceride to provide a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof.
- 30 40. A method of any one of claims 36 to 38, or a use of claim 39, wherein the pharmaceutical composition is stabilised to minimize the degradation of the ACE inhibitor or the pharmaceutically acceptable salt or derivative thereof.

41. A pharmaceutical composition, a method or a use of any one of the preceding claims, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is susceptible to heat and/or mechanical stress-induced degradation.

5

42. A pharmaceutical composition, a method or a use of claim 41, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril, trandolapril, quinapril, or a pharmaceutically acceptable salt or derivative thereof.

10

43. A pharmaceutical composition, a method or a use of claim 42, wherein the ACE inhibitor, or the pharmaceutically acceptable salt or derivative thereof, is ramipril or a pharmaceutically acceptable salt or derivative thereof.

15

44. A pharmaceutical composition, a method or a use of any one of the preceding claims, wherein the glyceride comprises one, two or three C₁₆-C₂₈ acyl moieties, wherein each C₁₆-C₂₈ acyl moiety is independently of the formula -CO-R, wherein R is a saturated or unsaturated hydrocarbon, which contains from 16 to 28 carbon atoms, and which is straight-chained or branched.

20

45. A pharmaceutical composition, a method or a use of claim 44, wherein R is a saturated hydrocarbon and/or wherein R is a straight-chained hydrocarbon.

25

46. A pharmaceutical composition, a method or a use of any one of the preceding claims, wherein the glyceride is a C₁₈-C₂₆ glyceride.

47. A pharmaceutical composition, a method or a use of claim 46, wherein the glyceride is a C₂₀-C₂₄ glyceride.

30

48. A pharmaceutical composition, a method or a use of any one of the preceding claims, wherein the glyceride comprises at least 50% diglyceride.

49. A pharmaceutical composition, a method or a use of any one of the preceding claims, wherein the glyceride is glycerol dibehenate.

Abstract

Pharmaceutical Composition

5 The present invention relates to a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof. In particular, the invention relates to a pharmaceutical composition, which comprises an ACE inhibitor, or a pharmaceutically acceptable salt or derivative thereof, and a C₁₆-C₂₈ glyceride. ACE inhibitors useful in the present invention are susceptible to
10 heat and/or mechanical stress-induced degradation. Preferred ACE inhibitors are ramipril, trandolapril, quinapril and pharmaceutically acceptable salts and derivatives thereof. The composition of the present invention may be for use as a medicament for the treatment or prevention of a cardiovascular disease, a coronary heart disease, a cerebrovascular disease, a peripheral vascular disease, arrhythmia, hypertension,
15 cardiac failure, cardiovascular death, myocardial infarction, stroke or angina.

The present invention further relates to a method of preparing the pharmaceutical composition of the present invention. The present invention also relates to a method of providing a stable pharmaceutical composition comprising an ACE
20 inhibitor, or a pharmaceutically acceptable salt or derivative thereof, by incorporating a C₁₆-C₂₈ glyceride into the composition. The present invention further relates to a use of a C₁₆-C₂₈ glyceride to provide a stable pharmaceutical composition comprising an ACE inhibitor or a pharmaceutically acceptable salt or derivative thereof.



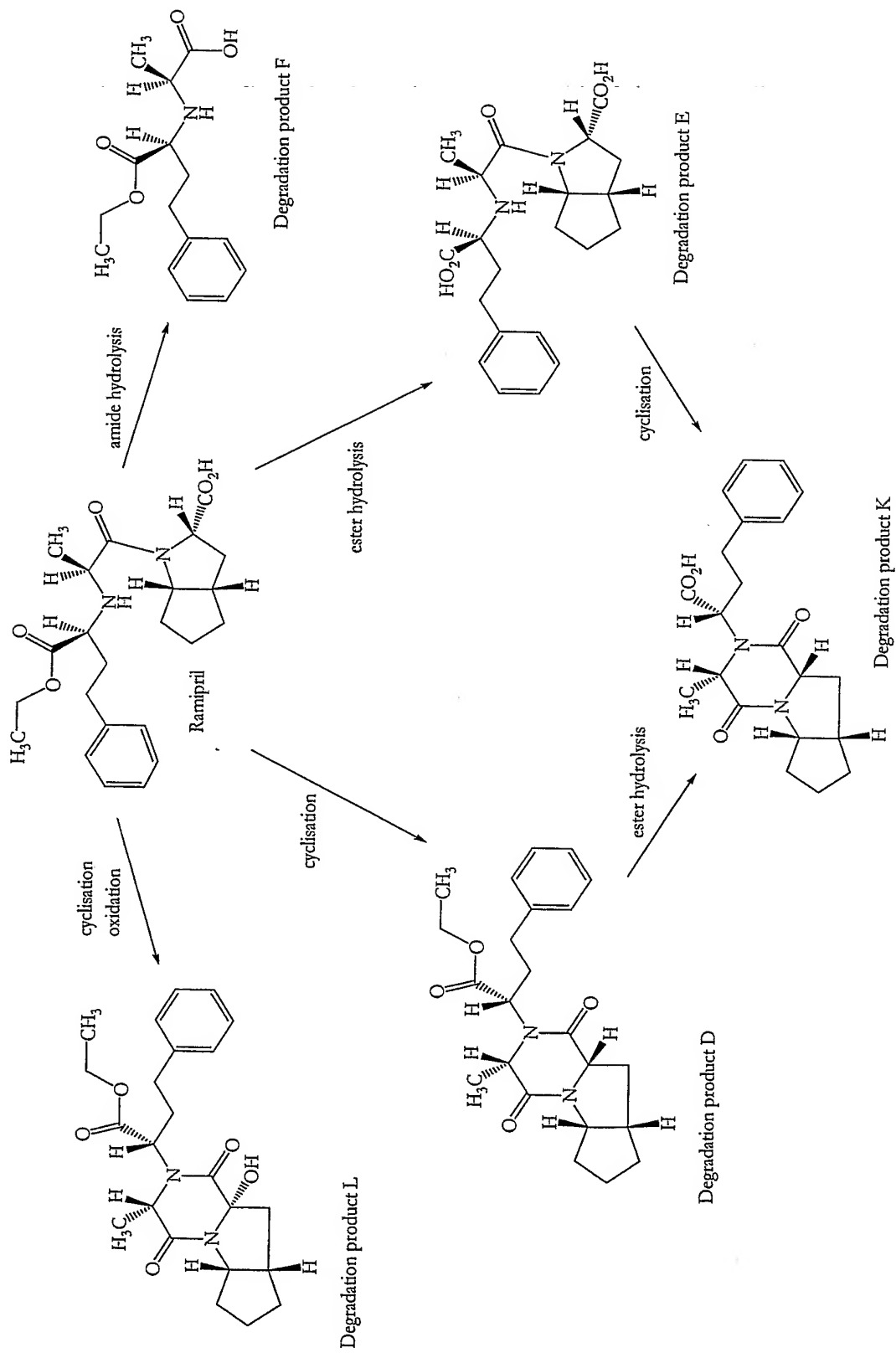


Figure 1



Total Impurities (%) at 40°C and 75% RH

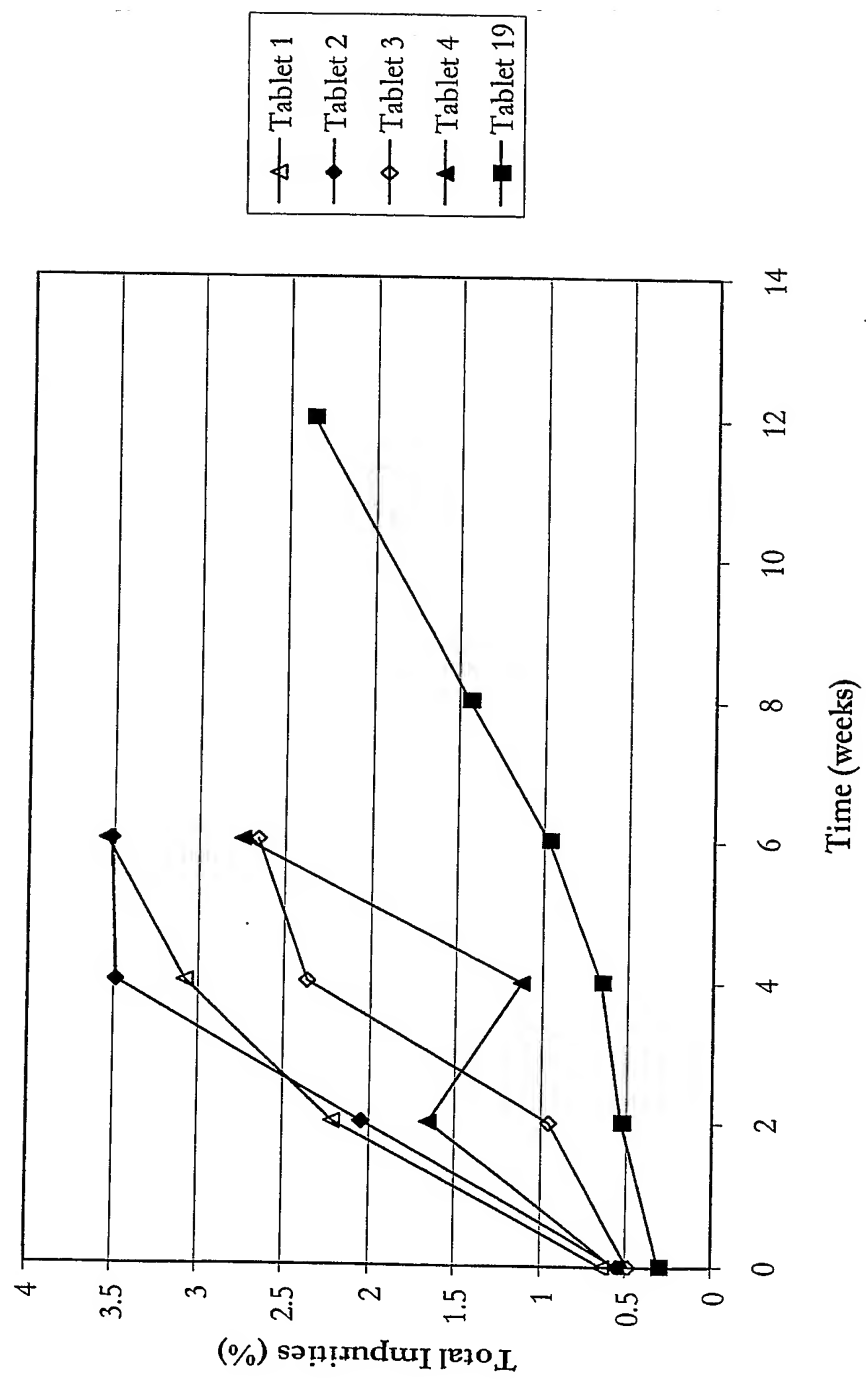


Figure 2

